APPLICATION NO. DATE

## INVENTOR SEARCH

=> d ibib abs 15 1-5

L5 ANSWER 1 OF 5 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER: 2008:978651 HCAPLUS Full-text

DOCUMENT NUMBER: 149:217060

TITLE: Modified flagellin with improved toll-like receptor 5

stimulating activity

INVENTOR(S): Rhee, Joon Haeng; Lee, Shee Eun;

Kim, Soo Young

PATENT ASSIGNEE(S): Chonnam National University, S. Korea

KIND DATE

SOURCE: PCT Int. Appl., 35pp.

CODEN: PIXXD2

LANGUAGE: English
FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

FAI	IX TIM	_	DAIE			VEEP	ICAI		DATE										
WO	2008	0970	A1		20080814			WO 2	008-		20080205								
	W: AE, AG, AL, A		AM,	AO,	AT,	AU,	ΑZ,	BA,	BB,	BG,	BH,	BR,	BW,	BY,	BZ,				
		CA,	CH,	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DO,	DZ,	EC,	EE,	EG,	ES,		
		FI,	GB,	GD,	GE,	GH,	GM,	GT,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,		
		KG,	KM,	KN,	KP,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LY,	MA,	MD,	ME,		
		MG,	MK,	MN,	MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,		
		PT,	RO,	RS,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	ΤJ,	TM,	TN,		
		TR,	TT,	TZ,	UA,	UG,	US,	UΖ,	VC,	VN,	ZA,	ZM,	zw						
	RW:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FΙ,	FR,	GB,	GR,	HR,	HU,		
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							CI,												
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			ΑZ,	ΒY,			MD,												
	8238				B1								20070209						
													20080204						
EP									EP 2008-712360 DK, EE, ES, FI, FR,										
	R:																		
				IT,	LI,	LT,	LU,	LV,	MC,	MT,	NL,	NO,	PL,	PT,	RO,	SE,	SI,		
	SK, TR																		
	CN 101622272						2010												
	IN 2009MN01473						2009	1211											
PRIORITY	PRIORITY APPLN. INFO.:									KR 2									
							KR 2008-11330						A 20080204 W 20080205						
										WO 2008-KR709 W 20080205							400		

AB Disclosed herein are flagellin mutants having an enhanced activity of stimulating the toll-like receptor-5 (hereinafter referred to as "TLR5"). The present invention relates to flagellin mutants, prepared by point-mutating some of the amino acids of a TLR5 agonist flagellin, such that flagellin mutantion suppress the multimerization of flagellin monomers, thus showing an enhanced activity of stimulating TLR5. The present inventor have prepared recombinant flagellin mutants, which can suppress polar-charge reactions that are involved in the axial interaction between flagellin monomers, of the flagellin gene flaB of V. vulnificus, by changing amino acid residues anticipated to be involved in the axial interaction, and have found that the prepared flagellin mutants have significantly enhanced TLR5-stimulating activity compared to that of prior (wild type) flagellin proteins. Improved flagellin vaccine adjuvants were developed by providing flagellin mutants

10/585.880 5/4/10

having enhanced TLR-stimulating activity compared to that of a prior flagellin, which was found to show a potent mucosal vaccine adjuvant effect by stimulating TLR5.

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 2 OF 5 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER: 2007:1191280 HCAPLUS Full-text

8

DOCUMENT NUMBER: 147:422950

TITLE: Site directed mutagenesis polypeptide essential for in

vivo expression of Vibrio vulnificus, and anti-vibrio

live vaccine comprising the same INVENTOR(S): Rhee, Joon Haeng; Lee, Shee Eun;

Kim, Soc Young; Kim, Choon Mee; Kim, Young Ran

Industry Foundation of Chonnam National University, S. PATENT ASSIGNEE(S):

SOURCE: Repub. Korean Kongkae Taeho Kongbo, No pp. given

CODEN: KRXXA7 DOCUMENT TYPE: Patent

LANGUAGE: Korean FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
KR 2007050206	A	20070515	KR 2005-107504	20051110
KR 807988	B1	20080228		
IORITY APPLN. INFO.:			KR 2005-107504	20051110

PRIORITY APPLN. INFO.:

A PyrH mutant which is an essential factor of Vibrio vulnificus in vivo is provided to be able to be used for developing vaccine and detecting and developing antibacterial materials. The site directed mutagenesis polypeptide is prepared by substituting arginine, which is a 62nd amino acid of an NTP binding site among a PyrH amino acid sequence. The protein sequence of PyrH gene has been provided. The anti-vibrio vulnificus live vaccine comprises a strain having the site directed mutagenesis polypeptide.

L5 ANSWER 3 OF 5 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER: 2007:636508 HCAPLUS Full-text DOCUMENT NUMBER: 147:230583

TITLE:

SOURCE:

CORPORATE SOURCE:

The pyrH gene of Vibrio vulnificus is an essential in

vivo survival factor

AUTHOR(S): Lee, Shee Eun; Kim, Soo Young;

Kim, Choon Mee; Kim, Mi-Kwang; Kim, Young Ran; Jeong, Kwangjoon; Ryu, Hwa-Ja; Lee, Youn Suhk; Chung, Sun

Sik; Choy, Hyon E.; Rhee, Joon Haeng

Clinical Vaccine R&D Center and Genome Research Center

for Enteropathogenic Bacteria, Chonnam National

University, Gwangju, 501-746, S. Korea

Infection and Immunity (2007), 75(6), 2795-2801

CODEN: INFIBR; ISSN: 0019-9567 PHRLISHER.

American Society for Microbiology

DOCUMENT TYPE: Journal LANGUAGE: English

The authors have suggested an important role of the pyrH gene during the infectious process of Vibrio vulnificus. Previously, the authors have identified 12 genes expressed preferentially during human infections by using in vivo-induced antigen technol. Among the in vivo-expressed genes, pyrH encodes UMP kinase catalyzing UMP phosphorvlation. Introduction of a deletion mutation to the pvrH gene was lethal to V. vulnificus, and an insertional mutant showed a high frequency 10/585.880 5/4/10

of curing. The authors constructed a site-directed mutant strain (R62H/D77N) on Arg-62 and Asp-77, both predicted to be involved in UMP binding, and characterized the R62H/D77N strain compared with the previously reported insertional mutant. The authors further investigated the essential role of the pyrH gene in the establishment of infection using the R62H/D77N strain. Cytotoxicity was decreased in the R62H/D77N strain, and the defect was restored by an in trans complementation. The i.p. 50% LD of the R62H/D77N strain increased by 26- and 238,000-fold in normal and iron-overloaded mice, resp. The growth of the R62H/D77N strain in 50% HeLa cell lysate, 100% human ascitic fluid, and 50% human serum was significantly retarded compared to that of the isogenic wild-type strain. The R62H/D77N mutant also had a critical defect in the ability to survive and replicate even in iron-overloaded mice. These results demonstrate that pyrH is essential for the in vivo survival and growth of V. vulnificus and should be an attractive new target for the development of antibacterial drugs and replication-controllable live attenuated vaccines. OS.CITING REF COUNT: 2 THERE ARE 2 CAPLUS RECORDS THAT CITE THIS RECORD

(2 CITINGS)

REFERENCE COUNT: 38 THERE ARE 38 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 4 OF 5 HCAPLUS COPYRIGHT 2010 ACS on STN ACCESSION NUMBER: 2006:23695 HCAPLUS Full-text

DOCUMENT NUMBER: 144:106237

TITLE: A bacterial Flagellin, Vibrio vulnificus FlaB, has a

strong mucosal adjuvant activity to induce protective immunity

AUTHOR(S): Lee, Shee Eun; Kim, Soo Young;

Jeong, Byung Chul; Kim, Young Ran; Bae, Soo Jang; Ahn, Ouk Seon; Lee, Je-Jung; Song, Ho-Chun; Kim, Jung Mogg;

Chov, Hvon E.; Chung, Sun Sik; Kweon, Mi-Na;

Rhee, Joon Haeno

Research Institute of Vibrio Infection and Genome CORPORATE SOURCE:

Research Center for Enteropathogenic Bacteria, Chonnam National University Medical School, Gwangju, 501-746,

S. Korea

Infection and Immunity (2006), 74(1), 694-702 SOURCE:

CODEN: INFIBR; ISSN: 0019-9567

PUBLISHER: American Society for Microbiology DOCUMENT TYPE: Journal

LANGUAGE: English

Flagellin, the structural component of flagellar filament in various locomotive bacteria, is the ligand for Toll-like receptor 5 (TLR5) of host cells. TLR stimulation by various pathogen-associated mol. patterns leads to activation of innate and subsequent adaptive immune responses. Therefore, TLR ligands are considered attractive adjuvant candidates in vaccine development. In this study, we show the highly potent mucosal adjuvant activity of a Vibrio vulnificus major flagellin (FlaB). Using an intranasal immunization mouse model, we observed that coadministration of the flagellin with tetanus toxoid (TT) induced significantly enhanced TT-specific IqA responses in both mucosal and systemic compartments and IGG responses in the systemic compartment. The mice immunized with TT plus FlaB were completely protected from systemic challenge with a 200+ min. LD of tetanus toxin. Radiolabeled FlaB administered into the nasal cavity readily reached the cervical lymph nodes and systemic circulation. FlaB bound directly to human TLR5 expressed on cultured epithelial cells and consequently induced NF-KB and interleukin-8 activation. Intranasally administered FlaB colocalized with CD11c as patches in putative dendritic cells and caused an increase in the number of TLR5expressing cells in cervical lymph nodes. These results indicate that flagellin would serve as an efficacious mucosal adjuvant inducing protective immune responses through TLR5 activation. OS.CITING REF COUNT:

41 THERE ARE 41 CAPLUS RECORDS THAT CITE THIS

RECORD (41 CITINGS)

REFERENCE COUNT: 39 THERE ARE 39 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L5 ANSWER 5 OF 5 HCAPLUS COPYRIGHT 2010 ACS on STN

ACCESSION NUMBER: 2005:696770 HCAPLUS Full-text
DOCUMENT NUMBER: 143:171319

TITLE: Mucosal vaccine adjuvants containing

bacterial flegellins derived from as an active

component Vibrio vulnificus, Salmonella typhimurium and Listeria monocytogenes

INVENTOR(S): Rhee, Joon Haeng; Lee, Shee Eun;

Kim, Soo Young

PATENT ASSIGNEE(S): Chonnam National University, S. Korea

.....

SOURCE: PCT Int. Appl., 21 pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

	KIND DATE																			
			A1 20050804						2005-											
		W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	AZ,	BA,	BE	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,		
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	D2	EC,	EE,	EG,	ES,	FI,	GB,	GD,		
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	EP	1708				AI	2006	1011		EP	2005-		20050112							
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				0.0							CN 2005-80002321 KR 2006-709082									
	KR 2007017300 KR 795839								RR 2006-709082						20060310					
	IN 2006MN00806				B1 20080117 A 20070330				IN 2006-MN806					20060710						
	US 20080069844										US 2007-585880									
PPTO						N.I	MI 20080320				KR 2004-1974									
PRIORITY APPLN. INFO.:											WO 2005-KR103									
													-							

ASSIGNMENT HISTORY FOR US PATENT AVAILABLE IN LSUS DISPLAY FORMAT

The present invention relates to mucosal vaccine adjuvants containing flagellins, the structural component of flagella, originated from Vibrio vulnificus, Salmonella typhimurium, and Listeria monocytogenes as an active component. The flagellin proteins are derived from flaA, flaB, flaC, flaD, flaB and flaF genes or their mutants. Protein sequences and DNA sequences of the flagellins and encoding genes are claimed but not presented.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RESULTS FROM SEARCHES IN REGISTRY, CAPLUS, MEDLINE, BIOSIS, EMBASE, AND DRUGU

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=> d que stat 114
        5503 SEA FILE=HCAPLUS ABB=ON ?VACCINE? AND ?MUCOS?
           59 SEA FILE=HCAPLUS ABB=ON L6 AND ?FLAGELL?
L7
L8
            6 SEA FILE=HCAPLUS ABB=ON L7 AND ?VIBRIO?(W)?VULNIFICUS?
L9
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L10
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            7 SEA FILE=HCAPLUS ABB=ON L8 OR L10
L11
L12
            13 SEA L11
L13
            12 DUP REMOV L11 L12 (8 DUPLICATES REMOVED)
T.14
            4 SEA L13 AND (PRD<20040112 OR PD<20040112)
L14 ANSWER 1 OF 4 HCAPLUS COPYRIGHT 2010 ACS on STN
ACCESSION NUMBER:
                       2003:490996 HCAPLUS Full-text
DOCUMENT NUMBER:
                        139:67779
TITLE:
                        Fusion proteins comprising an isolated pathogen
                        associated molecule pattern and an immunostimulatory
                       portion of an antigen for use as vaccines
INVENTOR(S):
                       Medzhitov, Ruslan; Kopp, Elizabeth
                      Yale University, USA
PATENT ASSIGNEE(S):
SOURCE:
                       PCT Int. Appl., 99 pp.
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Pat.ent.
LANGUAGE:
                        English
FAMILY ACC. NUM. COUNT: 3
PATENT INFORMATION:
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PATENT NO.						D	DATE			APPL	ICAT	ION I	NO.		DATE				
WO	WO 2003051305					A2 2003			30626 WO 2002-US40					20021213 <-					
WO	2003051305				A3 20040429														
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AU	2002	3616	82		A1		2003	0630		AU 2	002-	3616	82	20021213 <					
AU	AU 2007204086						2007	0830		AU 2007-204086					20070808 <				
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							AU 2001-286405						A3 20010731 <						
						WO 2	002-1	JS40	046		W 2	0021	213	<					

AB The present invention provides novel vaccines, methods for the production of such vaccines and methods of using such vaccines. The vaccines comprise chimeric protein of a pathogen associated mol. pattern (PAMP) and a antigenic epitope. The PAMPs are targets of innate immune recognition, e.g. chaperone, FimC; and the antigenic epitope is derived from pathogen antigen, tumor antigen, allergen, neural defect-related antigen, cardiovascular disease, rheumatoid arthritis-related antigen antigen, hormone, pregnancy-related antigen, embryonic antigen or fetal antigen. The novel vaccines of the present invention combine both of the signals necessary to activate native T-cells-a specific antigen and the co-stimulatory signal-leading to a robust and specific T-cell immune response. OS.CITING REF COUNT:

THERE ARE 1 CAPLUS RECORDS THAT CITE THIS RECORD

(1 CITINGS)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L14 ANSWER 2 OF 4 MEDLINE on STN

ACCESSION NUMBER: 2000323311 MEDLINE Full-text

DOCUMENT NUMBER: PubMed ID: 10862792

TITLE: WO 2003051305 flagellin that causes

IL-8 release from intestinal

epithelial cells.

AUTHOR: Steiner T S; Nataro J P; Poteet-Smith C E; Smith J A;

Guerrant R L

CORPORATE SOURCE: Division of Geographic and International Medicine,

University of Virginia Health Sciences Center,

Charlottesville, Virginia, USA.. ts5x@virginia.edu
CONTRACT NUMBER: AI-01573 (United States NIAID NIH HHS)

AI-26512 (United States NIAID NIH HHS)

AI-33096 (United States NIAID NIH HHS)

SOURCE: The Journal of clinical investigation, (2000 Jun)

Vol. 105, No. 12, pp. 1769-77.

Journal code: 7802877. ISSN: 0021-9738. L-ISSN: 0021-9738.

Report No.: NLM-PMC378507.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

(RESEARCH SUPPORT, NON-U.S. GOV'T) (RESEARCH SUPPORT, U.S. GOV'T, P.H.S.)

LANGUAGE: English

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 200007
ENTRY DATE: Entered STN: 10 Aug 2000

Last Updated on STN: 25 Jan 2002

Entered Medline: 24 Jul 2000

AB Enteroaggregative Escherichia coli (EAEC) is an emerging cause of acute and persistent diarrhea worldwide. EAEC infections are associated with intestinal inflammation and growth impairment in infected children, even in the absence of diarrhea. We previously reported that prototype EAEC strains rapidly induce IL-8 production by Caco-2 intestinal epithelial cells, and that this effect is mediated by a soluble, heat-stable factor released by these bacteria in culture. We herein report the cloning, sequencing, and expression of this biologically active IL-8-releasing factor from EAEC, and its identification as a flagellin that is unique among known expressed proteins. Flagella purified from EAEC 042 and several other EAEC isolates potently release IL-8 from Caco-2 cells: an engineered aflagellar mutant of 042 does not release IL-8. Finally, cloned EAEC flaggallin expressed in nonpathogenic E. coli as a polyhistidine-tagged fusion protein maintains its proinflammatory activity. These findings demonstrate a major new means by which EAEC may cause intestinal inflammation, persistent diarrhea, and growth impairment that characterize human infection with these organisms. Furthermore, they open new approaches for diagnosis and vaccine development. This novel pathogenic mechanism of EAEC extends an emerging paradigm of bacterial flagella as inflammatory stimuli.

L14 ANSWER 3 OF 4 MEDLINE on STN

ACCESSION NUMBER: 1996071810 MEDLINE Full-text

DOCUMENT NUMBER: PubMed ID: 7580302

TITLE: Synthetic recombinant vaccines against viral

agents.

AUTHOR: Arnon A; Levi R

10/585.880 5/4/10

CORPORATE SOURCE: Department of Chemical Immunology, Weizmann Institute of

Science, Rehovot, Israel.

International archives of allergy and immunology, SOURCE:

(1995 Dec) Vol. 108, No. 4, pp. 321-6. Ref: 48

Journal code: 9211652, ISSN: 1018-2438, L-ISSN: 1018-2438.

PUB. COUNTRY: Switzerland

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

General Review; (REVIEW)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199512

Entered STN: 24 Jan 1996 ENTRY DATE:

Last Updated on STN: 24 Jan 1996

Entered Medline: 19 Dec 1995

AB Synthetic recombinant vaccines are expression vectors incorporating defined epitope(s) of microbial agents. They are prepared by inserting synthetic oligonucleotide(s) coding for previously identified relevant epitopes into the genome of a desired vector, using recombinant DNA technology. The results discussed indicate that immunization with such vaccines carrying viral epitopes may lead to protective immunity against viral agents. Oligonucleotides coding for three influenza epitopes stimulating B cells, T helper cells and cytotoxic lymphocytes were individually inserted into the flagellin gene of a Salmonella vaccine strain. Immunization of mice with the resultant recombinant bacteria or their isolated flagella induced a specific mucosal anti-influenza protective response. The most efficient vaccine consisted of all three recombinant flagella, administered intranasally. The protection elicited was cross-strain specific, long-lasting and efficient

L14 ANSWER 4 OF 4 MEDLINE on STN

against a lethal viral challenge.

ACCESSION NUMBER: 1986165648 MEDLINE Full-text

DOCUMENT NUMBER: PubMed ID: 3514359

TITLE: Western blot analysis of intestinal secretory

immunoglobulin A response to Campylobacter jejuni antigens

in patients with naturally acquired Campylobacter

enteritis.

AUTHOR: Winsor D K Jr; Mathewson J J; DuPont H L

SOURCE: Gastroenterology, (1986 May) Vol. 90, No. 5 Pt 1,

pp. 1217-22.

Journal code: 0374630, ISSN: 0016-5085, L-ISSN: 0016-5085.

United States

PUB. COUNTRY: DOCUMENT TYPE: (COMPARATIVE STUDY)

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals

ENTRY MONTH: 198605

ENTRY DATE: Entered STN: 21 Mar 1990

Last Updated on STN: 21 Mar 1990

Entered Medline: 15 May 1986

AB Secretory immunoglobulin A (sIgA) response at the intestinal mucosa is a primary defense against enteric infections. We sought to determine which antigens of Campylobacter jejuni outer membranes elicited sIgA responses in 8 patients with naturally acquired Campylobacter enteritis using Western blot analysis of fecal extracts. Naturally acquired Campylobacter infection elicited an sIqA response in 7 of 8 patients. Of these 7 patients, 5 had Campylobacter-specific sIgA titers of 1:16 and two had titers of 1:64. The C. jejuni antigens eliciting sIgA production varied, but 5 of 8 patients exhibited reactions to a 63-kilodalton flagellar antigen, and 7 of 8 patients had a reaction with a 58- and a 44-kilodalton antigen of C. jejuni and

Campylobacter coli. Reaction with a 14.5- and a 97-kilodalton antigen was observed with the only stool that contained gross blood and mucus. Reactions with Campylobacter antigens were not detected in the fecal extracts of 5 healthy individuals. Identification of the antigens of C. jejuni that elicit an sIgA response may help us to better understand the immunology of Campylobacter enteritis and to identify antigens that are important in vaccine development.

## SEARCH HISTORY

# => d his ful

(FILE 'HOME' ENTERED AT 16:03:41 ON 04 MAY 2010)

FILE 'HCAPLUS' ENTERED AT 16:03:57 ON 04 MAY 2010

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L1 73 SEA ABB=ON ("RHEE JOON"/AU OR "RHEE JOON HAENG"/AU OR "RHEE JOON HANG"/AU)

E LEE SHEE EUN/AU

L2 41 SEA ABB=ON "LEE SHEE EUN"/AU

E KIM SOO YOUNG/AU

L3 256 SEA ABB=ON ("KIM SOO YOUN"/AU OR "KIM SOO YOUNG"/AU)

L4 19 SEA ABB=ON L1 AND L2 AND L3

L5 5 SEA ABB=ON L4 AND ?VACCINE?

FILE 'REGISTRY' ENTERED AT 16:21:22 ON 04 MAY 2010 E VIBRIO VULNIFICUS/CN

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FILE 'HCAPLUS' ENTERED AT 16:21:44 ON 04 MAY 2010

16 5503 SEA ABB=ON 2VACCINE? AND ?MUCOS?

17 59 SEA ABB=ON L6 AND ?FLAGELL?
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L8 6 SEA ABB=ON L7 AND ?VIBRIO?(W)?VULNIFICUS? L9 59 SEA ABB=ON L7 OR L8 L10 1 SEA ABB=ON L9 AND ?ISOLAT?(L)?BACT?

L11 7 SEA ABB=ON L8 OR L10

FILE 'MEDLINE, BIOSIS, EMBASE, DRUGU' ENTERED AT 16:23:14 ON 04 MAY 2010 L12 13 SEA ABB=ON L11

FILE 'HCAPLUS, MEDLINE, BIOSIS, EMBASE, DRUGU' ENTERED AT 16:23:52 ON 04 MAY 2010

L13 12 DUP REMOV L11 L12 (8 DUPLICATES REMOVED)

L14 4 SEA ABB=ON L13 AND (PRD<20040112 OR PD<20040112)

### FILE HOME

# FILE HCAPLUS

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FILE LAST UPDATED: 3 May 2010 (20100503/ED)
REVISED CLASS FIELDS (/NCL) LAST RELOADED: Feb 2010
USPTO MANUAL OF CLASSIFICATIONS THESAURUS ISSUE DATE: Feb 2010

 ${\tt HCAplus}$  now includes complete International Patent Classification (IPC) reclassification data for the first quarter of 2010.

CAS Information Use Policies apply and are available at:

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#### FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

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# http://www.cas.org/support/stngen/stndoc/properties.html

#### FILE MEDLINE

FILE LAST UPDATED: 2 May 2010 (20100502/UP). FILE COVERS 1947 TO DATE.

MEDLINE and LMEDLINE have been updated with the 2010 Medical Subject Headings (MeSH) vocabulary and tree numbers from the U.S. National Libra of Medicine (NLM). Additional information is available at

http://www.nlm.nih.gov/pubs/techbull/nd09/nd09\_medline\_data\_changes\_2010.

The Medline file has been reloaded effective January 24, 2010. See HELP RLOAD for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

See HELP RANGE before carrying out any RANGE search.

#### FILE BIOSIS

FILE COVERS 1926 TO DATE.

CAS REGISTRY NUMBERS AND CHEMICAL NAMES (CNs) PRESENT FROM JANUARY 1926 TO DATE.

RECORDS LAST ADDED: 28 April 2010 (20100428/ED)

BIOSIS has been augmented with 1.8 million archival records from 1926 through 1968. These records have been re-indexed to match current BIOSIS indexing.

#### FILE EMBASE

FILE COVERAGE: EMBASE-originated material 1974 to 4 May 2010 (20100504/ED Unique MEDLINE content 1948 to present

 ${\tt EMBASE}$  is now updated daily. SDI frequency remains weekly (default) and biweekly.

This file contains CAS Registry Numbers for easy and accurate substance identification.

For further assistance, please contact your local helpdesk.

## FILE DRUGU

FILE LAST UPDATED: 29 APR 2010 <20100429/UP>

>>> DERWENT DRUG FILE (SUBSCRIBER) <<<

>>> FILE COVERS 1983 TO DATE <<<

>>> THESAURUS AVAILABLE IN /CT <<<